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FA17

(54) [Title of the Invention] Electric deionizer

(57) [Abstract]

[Issue] To prevent a reduction in performance of an ion conductive substance which fills the electrode chambers of an electric deionizer, thereby enabling long-term operation.

[Means of Resolution] An electric deionizer comprising:

an anode chamber 17 having an anode 11; a cathode chamber 18 having a cathode 12; and enriching chambers 15 and desalination chambers 16 which are alternately formed by a plurality of anion exchange membranes 13 and cation exchange membranes 14 which are alternately arranged between the anode chamber 17 and cathode chamber 18. The anode chamber 17 and cathode chamber 18 are filled with an ion conductive substance, and deanionized water or pure water passes through the anode chamber 17, while de-cationized water or pure water 20 passes through the cathode chamber 18.

raw water

11: anode

12: cathode

25 13: A membrane

14: C membrane

15: enrichment chamber

16: desalination chamber

17: anode chamber

30 18: cathode chamber

waste water

enriched water

waste water

production water

[Scope of the Patent Claims]

[Claim 1] An electric deionizer comprising: an anode chamber having an anode; a cathode chamber having a cathode; and enriching chambers and desalination chambers which are alternately formed by a plurality of anion exchange membranes and cation exchange membranes which are alternately arranged between the anode chamber and cathode chamber; and

an enrichment chamber lies adjacent to the anode
10 chamber with a cation exchange membrane interposed, and
an enrichment chamber lies adjacent to the cathode
chamber with an anion exchange membrane interposed,
wherein the anode chamber and the cathode chamber are
filled with an ion conductive substance;

- de-anionized water or pure water passes through the anode chamber; and de-cationized water or pure water passes through the cathode chamber.
- 20 [Claim 2] The electric deionizer as claimed in claim 1, wherein the ion conductive substance in the anode chamber is an ion exchange resin and/or activated carbon, and the ion conductive substance in the cathode chamber is an ion exchange resin.
- [Claim 3] The electric deionizer as claimed in claim 2, wherein a heavy metal is supported by the ion conductive substance.
- 30 [Claim 4] The electric deionizer as claimed in any one of claims 1 to 3, wherein the cathode chamber outflow water is used as the anode chamber inflow water.

[Detailed Description of the Invention]

35 [0001]

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[Technical Field of the Invention] The present invention relates to an electric deionizer, and in particular the invention relates to an electric deionizer in which the electrical efficiency is

improved by filling the electrode chambers (anode chamber and cathode chamber) thereof with an ion conductive substance, and a reduction in the performance of the ion conductive substance which fills the electrode chambers of the electric deionizer is prevented, thereby enabling long-term operation.

[0002]

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[Prior Art] Electric deionizers such as that shown in 10 Figure 2, in which enrichment chambers 15 desalination chambers 16 are alternately formed by a plurality of anion exchange membranes (A membranes) 13 and cation exchange membranes (C membranes) 14 which are alternately arranged between electrodes (anode 11, 15 cathode 12), and in which the desalination chambers 16 are filled with a mixture or stacked layers of anion exchange material or cation exchange comprising ion exchange resins, ion exchange fibers or graft exchange material etc., are widely used in the 20 production of deionized water for use in various industrial and commercial fields and facilities, including semiconductor production plants, liquid crystal production plants, the pharmaceutical industry, the food industry, and the power industry, 25 among others (Japanese Patents 1782943, 2751090, 2699256). In Figure 2, 17 is an anode chamber and 18 is a cathode chamber, and spacers are generally provided.

[0003] Ions which flow into the desalination chambers
16 react with the ion exchange material due to the
affinity, concentration and mobility of these ions,
which then move through the ion exchange material in
the direction of the potential gradient, and move
across the membrane, and a neutral charge is maintained
35 in all of the chambers. The ions are decreased in the
desalination chambers 16 and are enriched in the
adjacent enrichment chambers 15 because of the
characteristics of the membranes which are selectively
permeable to ions, and because of the directivity of

the potential gradient. That is to say, cations pass through the cation exchange membranes 14, and anions pass through the anion exchange membranes 13 so as to be concentrated in the enrichment chambers 15. Consequently, deionized water (pure water) is recovered from the desalination chambers 16 as production water.

[0004] Raw water is introduced into the desalination chambers 16 and enrichment chambers 15 and deionized water (pure water) is recovered from the desalination chambers 16. Meanwhile, some of the enriched water which has been enriched with the ions flowing out from the enrichment chambers 15 is circulated to the inlet side of the enrichment chambers 15 by means of a pump (not depicted) in order to raise the water recovery rate, and the remainder is discharged to outside the system as waste water in order to prevent ion enrichment inside the system.

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20 [0005] Moreover, electrode water is also allowed to pass through the anode chamber 17 and cathode chamber 18, and this electrode water is replenished with conductive water having electrical conductivity of several tens of μS/cm or greater, or an NaCl 25 electrolyte or similar is added thereto in order to ensure conductivity.

[0006] Japanese Unexamined Patent Application Publication H10-43554 proposes filling the cathode 30 chamber with conductive particles, and USP 5,868,915 proposes filling the electrode chambers with an ion conductive substance; when the electrode chambers are filled with an ion conductive substance in this way, the ion conductive substance makes it possible to 35 ensure conductivity in the electrode chambers, and therefore it is no longer necessary to add the electrolyte to the electrode water or to replenish same with conductive water

[0007]

[Issues to be Resolved by the Invention] With electric deionizers in which the electrode chambers are filled with an ion conductive substance, there are advantages in that the electrical resistance is low and the electrical efficiency is high in the electrode chambers, but there are problems in that the ion conductive substance deteriorates because of oxidizing agents such as chlorine which are generated in the anode chamber; there are further problems in that there is a reduction in performance of the ion conductive substance because of scale deposition in the cathode chamber, and also in that operation cannot be continued over prolonged periods of time.

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[0008] The present invention aims to resolve these problems by providing an electric deionizer in which reductions in performance of the ion conductive substance which fills the electrode chambers are prevented, thereby enabling long-term operation.

[0009]

[Means of Resolving the Issues] The electric deionizer according to the present invention comprises: an anode 25 chamber having an anode; a cathode chamber having a cathode: and enriching chambers and desalination chambers which are alternately formed by a plurality of anion exchange membranes and cation exchange membranes which are alternately arranged between the anode chamber and cathode chamber; and an enrichment chamber lies adjacent to the anode chamber with a cation exchange membrane interposed, and an enrichment chamber lies adjacent to the cathode chamber with an anion exchange membrane interposed, wherein the anode chamber 35 and the cathode chamber are filled with an ion conductive substance; de-anionized water or pure water passes through the anode chamber; and de-cationized water or pure water passes through the cathode chamber.

[0010] The electrode chambers of the electric deionizer according to the present invention are filled with an ion conductive substance, and therefore the electrical resistance is low and the electrical efficiency is high in the electrode chambers.

[0011] In the cathode chamber of the electric deionizer, OH is generated by the reaction in (1) below. Then, when the electrode water which contains cations passes through the cathode chamber which is filled with the ion conductive substance, calcium scale is generated in the cathode chamber by the reaction in (2) below, and the calcium scale generating reaction is promoted by OH-.

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$$2H_2O + 2e^- \rightarrow 2OH^- + H_2$$
 ...(1)
 $Ca^{2^+} + CO_3^{2^-} \rightarrow CaCO_3$...(2)

[0012] Consequently, when water containing Ca2+ ions 20 passes through the cathode chamber which is filled with the ion conductive substance, there is a reduction in the performance of the ion conductive substance because of the scale which is generated in the cathode chamber.

25 [0013] In the present invention, de-cationized water, from which Ca^{2+} ions or the like have been removed, or pure water passes through the cathode chamber, and therefore scale is prevented from being generated and there is no reduction in performance of the ion 30

conductive substance.

[0014] Furthermore, the following reactions in (3) and (4) take place in the anode chamber, where chlorine is generated and the ion conductive substance such as an ion exchange resin undergoes oxidative degradation caused by the generation of chlorine.

$$2H_2O \rightarrow 4H^- + 4e^- + O_2$$
 ...(3)
 $2C1^- \rightarrow 2e^- + C1_2$...(4)

[0015] In the present invention, de-anionized water, from which Cl⁻ ions or the like have been removed, or pure water passes through the anode chamber, and therefore chlorine generation is prevented and oxidative degradation of the ion conductive substance is prevented.

[0016] Ion exchange resin may be cited as an ion conductive substance for filling the cathode chamber and the anode chamber, but ozone (03) which has a strong oxidizing power is also generated besides chlorine in the anode chamber, and therefore it is effective to use activated carbon which has a reducing action; the activated carbon may be used alone or ion exchange resin and activated carbon may be mixed together for use.

[0017] Furthermore, if a substance which supports a heavy metal is used as the ion conductive substance, it is possible to achieve an effect whereby the electrical resistance is further reduced because the heavy metal acts to increase conductivity, and this is preferred.

25 [0018] Furthermore, the cathode chamber outflow water contains hydrogen which is generated in the reaction (1) above. The cathode chamber outflow water containing this reducing hydrogen serves as the anode chamber inflow water, and this means that it is possible to reduce and remove the oxidizing substances generated in the anode chamber, and it is possible to more reliably prevent oxidative degradation of the ion conductive substance in the anode chamber. The cathode outflow water exits the cathode chamber and passes through the 35 anion exchange membranes whereby the anion component such as Cl is removed therefrom, which means that this water can be effectively utilized as water from which the anion component has been removed that passes through the anode chamber.

[0019]

[Mode of Embodiment of the Invention] A mode of embodiment of the present invention will be described below in detail with reference to the figures.

[0020] Figure 1 is a schematic view in cross section of the electric deionizer which represents a mode of embodiment of the present invention.

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[0021] Like the conventional electric deionizer shown in Figure 2, this electric deionizer is a device in which enrichment chambers 15 and desalination chambers 16 are alternately formed by a plurality of anion exchange membranes (A membranes) 13 and cation exchange membranes (C membranes) 14 which are alternately arranged between electrodes (anode 11, cathode 12), and in which the desalination chambers 16 are filled with a mixture or stacked layers of anion exchange material or 20 cation exchange material comprising ion exchange resins, ion exchange fibers or graft exchange material etc.

[0022] The anode chamber 17 and cathode chamber 18 are both adjacent to enrichment chambers 15, with a cation exchange membrane 14 and an anion exchange membrane 13 25 interposed, respectively, and the anode chamber 17 and cathode chamber 18 are filled with an ion conductive substance.

[0023] Raw water is introduced into the desalination 30 chambers 16 and enrichment chambers 15 and production water (pure water) is recovered from the desalination chambers 16. Some of the production water is pumped to the inlet side of the cathode chamber 18 as cathode 35 chamber 18 inflow water. The cathode chamber 18 outflow water is pumped to the inlet side of the anode chamber 17, and the anode chamber 17 outflow water is discharged to outside the system as waste water. Some of the enriched water which has been enriched with the ions flowing out from the enrichment chambers 15 is circulated to the inlet side of the enrichment chambers 15, and the remainder is discharged to outside the system as waste water.

[0024] With this electric deionizer, the anode chamber 17 and cathode chamber 18 are filled with the ion conductive substance, and therefore the electrical resistance is low and the electrical efficiency is high

1.0 in the electrode chambers.

[0025] Ion exchange material such as ion exchange resin, ion exchange fibers or graft exchange material may be cited as ion conductive substances; the ion conductive 1.5 substance used in the cathode chamber 18 is preferably a mixed ion exchange material such as a mixed bed ion exchange resin comprising anion exchange resin and cation exchange resin, or an anion exchange material alone such as anion exchange resin.

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2.5

[0026] On the other hand, the ion conductive substance used in the anode chamber 17 is preferably a mixed ion exchange material such as a mixed bed ion exchange resin comprising anion exchange resin and cation exchange resin, or a cation exchange material alone such as cation exchange resin, or a mixture of activated carbon and these ion exchange materials.

[0027] Filling the anode chamber 17 with activated carbon makes it possible to 30 prevent oxidative degradation in the anode chamber 17, by virtue of the reducing action of the activated carbon, described above, and this is preferable.

35 [0028] Furthermore, one or two or more types of heavy metals such as palladium, iron and manganese may be supported by some or all of the ion exchange materials such as the ion exchange resins which fill the cathode chamber 18 and anode chamber 17; when such heavy metals

are supported, an effect is demonstrated whereby the electrical resistance is further reduced. In this case, no particular limitation is imposed on the amount of heavy metal supported, but this amount is preferably around 0.5 - 10 wt% with respect to the ion exchange material.

[0029] Furthermore, when the ion exchange materials are used in conjunction with activated carbon, and when the ion exchange materials support heavy metals, the electrode chambers are filled with the ion exchange material which supports activated carbon or heavy metals along the surface of the electrode plates and this is effective in favoring reaction.

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[0030] With the electric deionizer in Figure 1, production water (pure water) flows through the cathode chamber 18 which is filled with the ion conductive substance in this way, and therefore scale is prevented from being generated in the cathode chamber 18, and so a reduction in the performance of the ion conductive substance caused by the scale is prevented.

[0031] Furthermore, the cathode chamber 18 outflow
25 water which flows through the anode chamber 17 is pure
water flowing into the anode chamber 17, and it does
not contain an anion component, and moreover it
contains reducing hydrogen which is generated in the
electrode reaction inside the cathode chamber 18, and
30 therefore it is possible to prevent oxidative
degradation of the ion conductive substance in the
anode chamber 17.

[0032] Moreover, the electric deionizer shown in Figure 1 is one example of a mode of embodiment of the electric deionizer according to the present invention, but the present invention is not limited to the device shown in the figure, provided that any variations remain within the scope of the invention.

[0033] For example, with the electric deionizer shown in Figure 1, the enrichment chambers 15 are not filled with the ion conductive substance, but the enrichment chambers may equally be filled with an ion exchange material or an ion conductive substance such activated carbon or heavy metal. Furthermore, with the electric deionizer shown in Figure 1, production water serves as the cathode chamber 18 inflow water, but pure 10 water from a separate system or water from which the cation component has been removed by processing raw water in a cation exchange resin column (soft water) may equally be used as the cathode chamber 18 inflow water. Furthermore, instead of using the cathode 15 chamber 18 outflow water for the anode chamber 17 inflow water, it is feasible to use production water or pure water from a separate system or water from which the anion component has been removed by processing raw water in an anion exchange resin column. It is not necessarily the case that some of the enriched water 20 has to be circulated, and when the quality of the production water is more of a priority than the water recovery rate, raw water may be allowed to flow through for a time in any direction.

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[0034]

[Effects of the Invention] As described above, the present invention provides an electric deionizer in which the electrical efficiency is improved by filling the electrode chambers thereof with an ion conductive substance, and a reduction in the performance of the ion conductive substance which fills the electrode chambers of the electric deionizer can be prevented, thereby enabling long-term operation.

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[Brief Description of the Figures]

[Figure 1] is a schematic view in cross section of an electric deionizer which represents a mode of embodiment of the present invention; and

[Figure 2] is a schematic view in cross section which represents a conventional electric deionizer.

[Key to Symbols]

- 5 11 anode
 - 12 cathode
 - 13 anion exchange membrane (A membrane)
 - 14 cation exchange membrane (C membrane)
 - 15 enrichment chamber
- 10 16 desalination chamber
 - 17 anode chamber
 - 18 cathode chamber

[Figure 1]

- 15 raw water
 - 11: anode
 - 12: cathode
 - 13: A membrane
 - 14: C membrane
- 20 15: enrichment chamber
 - 16: desalination chamber
 - 17: anode chamber
 - 18: cathode chamber

waste water

25 enriched water

waste water

production water

[Figure 2]

30 raw water

electrode water

13: A membrane

14: C membrane

electrode water

35 11: anode

12: cathode

15: enrichment chamber

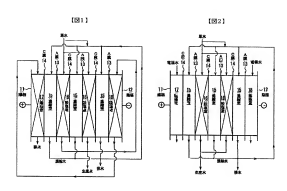
16: desalination chamber

17: anode chamber

18: cathode chamber enriched water

production water

waste water



PATENT ABSTRACTS OF JAPAN

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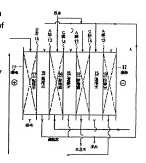
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(54) ELECTRIC DEIONIZATION EQUIPMENT

(57)Abstract:

PROBLEM TO BE SOLVED: To enable to operate for a long time by preventing the performance deterioration of an ion conductive substance filled in the electrode chamber of an electric deionization equipment is an equipment which is provided with an anode chamber 17 having an anode 11, a cathode chamber 18 having a cathode 12 and an enriching chamber 15 and desalination chambers 16 alternately formed by alternately disposing a plurality of anion exchange membranes and cation exchange membranes 14 between the chamber 17 and the chamber 18 are filled in the ion conductive substance, the water after an anion is



removed or a pure water is allowed to pass through the chamber 17 and the water after a cation is removed or the pure water is allowed to pass through the chamber 18.

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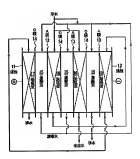
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(54) [発明の名称] 能気脱イオン装帽

(57)【要約】

【課題】 電気脱イオン装置の電極室に充填したイオン 導電性物質の性能低下を防止して長期運転を可能とす

【解決手段】 陽極11を有する陽極室17と、陰極1 2を有する除極室18と、これらの陽極空17と除極室 18との間に複数のアニオン交換膜13及びカチオン交 換購14を交互に配列することにより交互に形成された 濃縮室15及び劉塩室16とを備える電気脱イオン装 置。陽極変17及び陰極室18にイオン導電性物質が充 填されており、陽極変17にはアニオンを除去した水又 は純水が通水され、陰極室18にはカチオンを除去した 水又は純水が適水される。



【特許請求の範囲】

【請求項 1 】 陽極を有する陽極変と、陰極を有する陰極変と、これらの陽極変と陰極変との間に複数のアニオン交換膜及びカテオン交換膜及びカテオン交換膜を交互に配列することによ

り交互に形成された法確室及び脱塩室とを備え、 陽極室にはカテオン交換機を介して造確室が解接し、除 極室にはアニオン交換膜を介して造確室が解接している

電気脱イオン装置において. 陽極室及び陰極室にイオン導電性物質が充填されてお

陽極室にはアニオンを除去した水又は純水が通水され、 陰極室にはカチオンを除去した水又は純水が通水される ことを特徴とする電気脱イオン装置。

[論求項2] 結束項1でおいて、陽極室のイオン導電 性物質がイオン交換制階及び/又は活性膜であり、降極 室のイオン導電性物質がイオン交換機能であることを特 数とする電気脱イオン等麗。

[請求項3] 請求項2において、イオン練器性物質に 意金膜が指待されていることを特徴とする選気脱イオン 装置。 (請求項4] 請求項1ないし3のいずれか1項におい

て、除極壺流出水を陽極室流入水とすることを特徴とす る電気脱イオン鉄艦。

【発明の詳細な説明】

[0001]

(発野の属する政治分野) 会専明は電気組みオン経歴に 係り、谷に電気製みオン終更の電指室(特殊重要の経ង 恵) にイオン郷電性物質を売抜することにより電気効率 を高めた電気能イオン終重において、電極空に売損した イオン郷電性物理の性能能下を防止して長期環転を可能 30 とした電気組みオン終歴に防ちる。

[0002]

【役本の状態】②※、半高体製造工場、液晶製造工場、 製菓工業、食品工業、電力工業等の各種の企業欠及政性 用ないし研究組織等において使用される配イオンを収 造化は、国2と不予如く、電管(保施 11、降価 12) の間に開放のアニオン交換額(Aip)13及びカチオン 交換額(Cip)14を交互に応列して健康宣16とより 2 大機・イン交換額はもしくはグラフト交換件等からなる 現上、イオン交換額はもしくはグラフト交換件等からなる アニオン交換を減した電気担イオン表数が多用されている「俗辞 第17829439、特許第27510909、特許第 2699256号)。なお、国2において、17は解析 室、18は終格室であり、一般にスペー学が懲げられて いる。

[0003] 新塩室16に流入したイオンはその機和 力、造度及び移動度に基いてイオン交換体を反応し、電 塩の総もの方向にイオン交換体を原応し、環 増かて、陽極窒化なアンボータを開発したのでは 切って移動し、すべての窓におして電荷の中枢が保たれ 59 水が温水され、陰極窒にはアニオンを除去した水叉は純 が助って移動し、すべての窓におして電荷の中枢が保たれ 50 水が温水され、陰極窒にはアニオンを除去した水叉は純

る。そして、隣のイオン連邦的透過特性のため、及び電 位の階級の方向性のために、イオン減率機変16では減 少し、腕りの結確変15では減縮される。即ち、カキンはカオオン交換機14を透過して、また、アニオンは アニオン交換機13を透過して、それぞれ減縮至15内 に結婚される。このため、緊縄変16から生産水として 版イオン米(場外)が回収される。

[0004] 原状は沿極室 [6と議職室]5とに導入さ ・ 影塊室]5から比較、ナンボ (後外)が取り出され 10 る。一方、議職室 15から流出するイオンが議職された 通総材は、ポンプ (国元セギ)により一部が小国収率の 向上のために領庫宣 15の入間側に関係され、 内のイオンの活備を防止するために排水として系列へ横 置きれる。

[0005] なお、時程室17及び除極空18にも電極 水が通波されており、この電極水には、爆発性の発尿の ために数十 x x c m以上の電気伝導率を有する等電性 のある水を領給するか、或いはNaC1等の電解質が添 加される。

20 [0006] 特別平10-43554号公領には降極空 に電気伝導性子を充領するととが提案されており、ま た、USP5、868、915には、暗極空にイオン等 強性物質を充実することが提案されており、でのように 報燈変にイメン等機能物を充填した場合には、このイ オン等尾性物理により、電板空の博客性を確保すること ができるため、電径水への電影質の添加や場尾性の水の 補助を不受とすることができるため。

100071

【発酵が解決しようとする課題】 電極室にイオン場電性 物質を売壊した電気限イオン砂温は、電管室の電気拡充 か小さく、電気が建か高いという利点を有するが、時極 空では発生する埋荒寺の酸化添れよるイオン棒電性物質 の劣化の配配があり、また、 路極空ではスケール作出に よるイオン郷電性物質の性能低下の問題があり、 長寿温 転を継続するとができないという問題があり、 長寿温

[0008]本発明は上記従来の問題点を解決し、選性 変に実達したイオン構築性物質の性能低下を防止して表 制運転を可能とした鑑気貼イオン装置を提供することを 目的とする。

[0009]

[関連を解析するための手段] 本発明の電気脱イオン族 鑑認、際極を前する時極度と、特権を有する時極度と、 たちの特権型と機構を立ている。 及びおオナン炭原規を交互に基列するととより交互 板はおれた場合数とが回復型とから、 オン交換規を介して減極型の解析し、特権型にはフェオ ン受除規を介して減極型の解析し、存権型にはフェオ ン受除規を介して減極型の解析し、存権型にはフェオ としている。 原性型にはフェオ としている。 原性型にはフェオ としている。 のでは、 の 水が適水されることを特徴とする。

【0010】本発明の電気脱イオン装置は、電極室にイ オン導電性物質を充填したため、電極室の電気抵抗が小 さく、電気効率が高い。

【3011】電気脱イオン鉄置の陰極室においては、以 下の (1) の反応でOH が発生する。そして、カチオ ンを含有する電極水をイオン導電性物質が充填された除 極室に通水すると、陰極室においては、下記(2)の反 応でカルシウムスケールが生成し、このカルシウムスケ

ールの生成反応がOH* により促進される。 2H, O+2e - →2OH +H;

Ca2 + + CO, 2 - → CaCO. ... (2) 【0012】このため、Ca2・イオンを含む水をイオ

ン郷電性物質が充填された除極空に過水すると、除極窒 で発生するスケールのためにイオン導電性物質の性能低 下が生じる。

【0013】本発明では、Ca2+イオン等のカチオン を除去した水又は純水を除極室に通水するため. このよ うなスケールの生成が防止され、イオン導電性物質の性 能低下を生じることがない。

[0014]また、隣接室においては、以下の(3)。 (4)の反応が超とり、塩素が生成し、生成した塩素に よりイオン交換樹脂等のイオン導電性物質が酸化劣化す

2 H2 O-4 H + 4 e + O2 ... (3)

【0015】本発明では、C1 イオン等のアニオンを 除去した水又は純水を隔極室に通水するため、塩素の生 成が防止され、イオン導電性物質の酸化劣化が防止され

【0016】除極壺及び陽極壺に充填するイオン導電性 物質としてはイオン交換樹脂が挙げられるが、陽極壺に おいては、塩素以外にも酸化力の強いオゾン(〇。)が 発生するため、還元作用のある活性炭を用いることが効 果的であり、活性炭を単独で用いるか吸いは活性炭とイ オン交換樹脂とを混合して用いても良い。

【0017】また、イオン郷電性物質として、重金属を 担持したものを用いることにより、重金属が導電性を増 加させるという作用により、更に電気抵抗を低減させる という効果を得ることができ、好ましい。

[0018] また、陰極室の淺出水には、前記(1)の 反応で発生した水素が含まれている。この還元性の水素 を含む陰径空流出水を陽極室の流入水とすることによ り、陽極室で発生する酸化性物質を還元して除去するこ とができ、隔極室におけるイオン導電性物質の酸化劣化 をより確実に防止することができる。この陰極室流出水 は、陰極室からアニオン交換膜を透過してCI 等のア ニオン成分が除去されているため、陽頻室に通水するア ニオン成分除去木として有効に利用することができる。

[0019]

... (4) 2C1 - → 2e + Cl2

換樹脂、イオン交換繊維、グラフト交換体等のイオン交

30 換体が挙げられるが、陰極窒18のイオン導電性物質と しては、アニオン交換御脂とカチオン交換御脂との混床 イオン交換樹脂等の混合イオン交換体、或いはアニオン 交換樹脂等のアニオン交換体単独を用いるのが好まし

[0026] 一方、隣極室17のイオン導電性物質とし ては、アニオン交換制脂とカチオン交換制脂との縄床イ オン交換樹脂等の複合イオン交換体、或いはカチオン交 袋樹脂等のカチオン交換体単独、或いは活性炭、或いは 活性炭とこれらのイオン交換体との拠合物を用いること

40 ができる。 【9027】陽極室17に活性炭を充填することは、前 述の如く、活性炭の還元作用で陽極室 1.7 における酸化 出化を防止することができ、好ましい。

【9028】また、陰極室18及び陽極壺17に充填す るイオン交換樹脂等のイオン交換体の一部又は全部にパ ラジウム、鉄、マンガン等の重金属の1種又は2種以上 を狙持させても良く、このような重金属を担持させるこ とにより、更に電気抵抗を低減させるという効果が奏さ

れる。この場合、重金層の銀持量には特に制限はない 50 が、イオン交換体に対してり、5~10重量%程度とす

【発明の実施の形態】以下に図面を参照して本発明の実 飾の影響を詳細に説明する。

【0020】図1は本発明の実施の形態を示す電気脱イ

オン鉄圏の模式的な断面図である。

【0021】との電気脱イオン装置は、図2に示す従来 の電気脱イオン装置と同様、電極(陽極11、除極! 2)の間に複数のアニオン交換膜(A購) 13及びカチ オン交換膜 (C購) 1.4を交互に配列して濃縮室15と 脱塩室16とを交互に形成したものであり、脱塩室16

19 には、イオン交換樹脂、イオン交換機能もしくはグラフ ▶交換体等からなるアニオン交換体及びカチオン交換体 が混合もしくは複層状に充填されている。

[0 0 2 2] 陽極室 1 7及び終極室 1 8は各々カチオン 交換鎖14及びアニオン交換鎖13を介して濃縮室15 と隣接しており、この稀極室17及び陰極室18にはイ

オン郷電性物質が充填されている。 【0023】原水は脱塩窒16と濃縮窒15に導入さ

れ、脱塩空】6からは生産水 (純水) が取り出される。 この生産水の一部は、降極室18歳入水として降極室1 20 8の入口側へ送給される。陰極室18の流出水は陽極室 17の入口側へ送給され、陽極壺17の織出水は排水と して系外へ排出される。波磁室 1 5 から流出するイオン が濃縮された濃縮水は、一部が濃縮室 15 の入口側に循

躁され、残鄙は排水として系外へ排出される。 【0024】この電気脱イオン装置では、陽極室17及 び除極空18にイオン導電性物質が充填されているた め、電極室の電気抵抗が小さく、電気効率が高い。 [0025] このイオン導端性物質としては、イオン交 るのが好ましい。

[0029]また、イオン交換体と共に活性炭を併用したり、イオン交換体に重金膜を担待させる場合。活性炭や重金原担待イオン交換体と電程整内の電極板面に沿って索填し、優先的に反応させることも効果的である。

[0030] 図1の電気脱イオン修置では、このように イオン海産性頻度を充填した路極変18に生産水(稀水)を連水するため、陰極変18におけるスケール発生 が防止され、スケールによるイオン準電性物質の性能低 下が防止される。

[0031]また、特殊監2 7 に通水される総数室18 の走出水は、物水が特殊室17 に通水されたものであり、アエオルの入を含まず、しかも、保証室18 内での電販反応で生成した還元性物質の水素を含むため、特殊室17 におけるイオン等電性物質の取化素化が初止される。

[0032]なお、図1に示す電気脱イオン装置は本発明の電気脱イオン装置の実施の形態の一例であって、本発明はその製造を超えない限り、何ら図示のものに限定されるものではない。

[0033] 明えば、回1の電気限々オン装置では減額 直18はイメン環盤物質が売減されていないが、この 濃縮室に6イオン交換体、活性規又は金貨等のイオン準 電性物質が充填されていても良い。また、同りの電気限 イオン保護では速か水を構造。18の成入水をしている が、降極電18の使入水としては、別系板の熱水似は 原本をラチオン交換指数で必要があれている分粉を末 原本をラチオン交換指数で必要がある。

*水(秋水)を用いても良い。また、陽極重17の流入水 についても、陽極重88 88 元出水を用いる他、生産水や 駅系数の株火板いは原水をアンゴン交換制脂がで処理し たアニオン成分除去水を用いても良い。更に、濃硬水に ついても必ずしも一部を循環する必要はなく、水調収率 よりも生産水の水質が重複される場合には、原水を一通 性で流水しても良く、その流水方向等も任意である。

[0034]

[発明の効果]以上部途した遭り、本発明によれば、塞 10 種弦にイオン等面性物質を充填して電気効率を高めた電 気料イオン禁硬において、電極変に充填したイオン場電 性物質の性能能下を防止することができ、これにより長 筋送線症が5円能な電気料イオン装置が提供される。 [図面の物理と説明]

【図1】 本発明の実施の形態を示す電気腕イオン装置の 絶式的な筋面図である。

【図2】 従来の電気脱イオン装置を示す模式的な断面図 である。

【符号の説明】

20 11 陽極

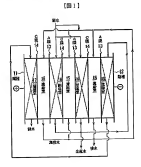
12 降極

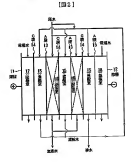
アニオン交換膜(A瞬)
 カチオン交換膜(C膜)

15 濃箱室

16 脱塩変

17 陽極空 18 陰極空





フロントページの続き

F ターム(参考) 4D006 GA17 1A30C KA31 KD19 MA13 MA14 PA01 P802 4D061 DA01 DB18 EA09 EB13 FA08 FA17